This Page Is Inserted by IFW Operations and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

As rescanning documents will not correct images, please do not report the images to the Image Problem Mailbox.



Europäisches Patentamt

European Patent Office

Office européen des brevets



(11)

EP 1 065 169 A1

(12)

EUROPEAN PATENT APPLICATION

- (43) Date of publication: 03.01.2001 Bulletin 2001/01
- (21) Application number: 00305433.5
- (22) Date of filing: 28.06.2000

- (51) Int. Cl.⁷: **C01G 23/047**, C01G 23/053, B01J 21/06, B01J 35/00
- (84) Designated Contracting States:
 AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU
 MC NL PT SE
 Designated Extension States:
 AL LT LV MK RO SI
- (30) Priority: 30.06.1999 JP 18573999 07.10.1999 JP 28681699
- (71) Applicant: Sumitomo Chemical Company, Limited Chuo-ku Osaka 541-8550 (JP)

- (72) Inventors:
 - Sakatani, Yoshiaki
 Niihama-shi, Ehime (JP)
 - Kolke, Hironobu Niihama-shi, Ehime (JP)
 - Takeuchi, Yoshlaki
 Nilhama-shi, Ehime (JP)
- (74) Representative:
 Cresswell, Thomas Anthony
 J.A. KEMP & CO.
 14 South Square
 Gray's Inn
 London WC1R 5LX (GB)

(54) Titanium oxide, photocatalyst comprising same and photocatalytic coating agent

(57) Titanium oxide having an index X_1 calculated by the following equation (I):

$$X_1 = \dot{B}_1/A_1 \tag{i}$$

of not more than about 0.90, and an index Y_1 calculated by the following equation (ii):

$$Y_1 = D_1/C_1$$
 (II)

of not less than about 0.075, wherein A_1 and B_1 represent respective half-widths of peaks, which are obtained by the process consisting of the steps:

- (i) analyzing titanlum oxide eight times according to an X-ray photoelectron spectroscopy,
- (ii) obtaining an integrated spectrum of an electron state of titanium with respect to the above first analysis and the second analysis,
- (iii) obtaining a half-width, A₁, of a peak within a binding energy range of from 458 eV to 460 eV with respect to the integrated spectrum obtained in the above step (ii), and
- (iv) carrying out the same steps as the above steps (ii) and (iii) with respect to the above seventh analysis and the eighth analysis to obtain a half-width, B₁, of a peak,
 - C₁ represents the integrated value of absorb-

ance within a wavelength range of from 250 nm to 550 nm in measurement of an ultraviolet-vislble diffuse reflection spectrum of titanium oxlde, and

D₁ represents the integrated value of absorbance of titanium oxide within a wavelength range of from 400 nm to 550 nm, is useful as a photocatalyst.



Description

.10

35

45

50

55

Field of the Invention

[0001] The present invention relates to titanium oxide, which exhibits a superior photocatalytic activity under irradiation of not only an ultraviolet light but also a visible light; a photocatalyst comprising said titantum oxide as a catalyst component; and a photocatalytic coating agent comprising said photocatalyst and a solvent.

Background of the Invention

Titanium oxide is known as a substance exhibiting an oxidation activity or reduction activity under light irra-[0002] diation. Hereinafter, such a substance is referred to as a photocatalyst. A photocatalyst containing said titanium oxide is available commercially. When such a photocatalyst is contacted with, for example, bad-smelling substances present in a living space and working space, or undesired substances such as organic solvents, agricultural chemicals and sur-15 face active agents present in water, said bad-smelling substances or undesired substances can be decomposed.

However, said commercially available photocatalyst cannot exhibit a superior photocatalytic activity under visible light irradiation, although it can exhibit a superior photocatalytic activity under ultraviolet light irradiation.

Summary of the Invention

[0004] Accordingly, an object of the present invention is to provide titanium oxide, which exhibits a superior photocatalytic activity under irradiation of visible light as well as ultraviolet light.

Another object of the present invention is to provide a photocatalyst comprising said titanium oxide as a catalyst component.

25 [0006] A further object of the present invention is to provide a photocatalytic coating agent comprising said photocatalyst and a solvent.

The present inventors have undertaken extensive studies about a photocatalyst. As a result, the present inventors have found a specific titanium oxide, which exhibits a superior photocatalytic activity under irradiation of visibie light as well as ultraviolet light. And thereby the present invention has been obtained.

The present invention provides titanium oxide having a value of an index X_1 calculated by the following equation () of not more than about 0.90, and a value of an index Y₁ calculated by the following equation (II) of not less than

$$X_1 = B_1/A_1 \tag{1}$$

$$Y_1 = D_1/C_1 \tag{II}$$

wherein, as to the equation (I), A₁ and B₁ stand for respective half-widths of peaks, which are obtained by the process

(i) analyzing titanium oxide eight times according to an X-ray photoelectron spectroscopy,

(ii) obtaining an integrated spectrum of an electron state of titanium with respect to the above first analysis and the

(iii) obtaining a half-width, A₁, of a peak within a binding energy range of from 458 eV to 460 eV with respect to the integrated spectrum obtained in the above step (ii), and

(iv) carrying out the same steps as the above steps (ii) and (iii) with respect to the above seventh analysis and the eighth analysis to obtain a half-width, B_{1} , of a peak, and

as to the equation (II), C₁ stands for an integrated value of absorbance within a wavelength range of from 250 nm to 550 nm in measurement of an ultraviolet-visible diffuse reflection spectrum of thankum oxide, and D_1 stands for an integrated value of absorbance of titanium oxide within a wavelength range of from 400 nm to 550 nm.

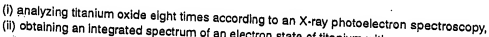
[00091 Further, the present invention provides titanium oxide having a value of an index X₁ calculated by the following equation (i) of not more than about 0.90, a value of an index Y1 calculated by the following equation (ii) of not less than about 0.075, and a value of an index Z₁ calculated by the following equation (III) of not less than about 0.75,

$$X_1 = B_1/A_1 \tag{I}$$

$$Y_1 = D_1/C_1 \tag{II}$$

$$Z_1 = Y_1 \times E_1$$

(III)wherein, as to the equation (I), A_1 and B_1 stand for respective half-widths of peaks, which are obtained by the process



10

15

25

30

35

40

45

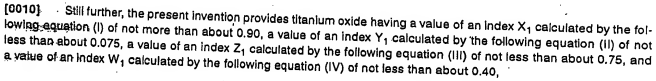
50

55

(ii) obtaining an integrated spectrum of an electron state of titanium with respect to the above first analysis and the

(iii) obtaining a half-width, A1, of a peak within a binding energy range of from 458 eV to 460 eV with respect to the integrated spectrum obtained in the above step (ii), and

(iv) carrying out the same steps as the above steps (ii) and (iii) with respect to the above seventh analysis and the eighth analysis to obtain a half-width, B_1 , of a peak; as to the equation (II), C_1 stands for an integrated value of absorbance within a wavelength range of from 250 nm to 550 nm in measurement of an ultraviolet-visible diffuse reflection spectrum of titanium oxide, and D₁ stands for an integrated value of absorbance of titanium oxide within a wavelength range of from 400 nm to 550 nm; and as to the equation (III), E₁ stands for a crystallite size of titanium Section Section



$$X_1 = B_1/A_1 \tag{I}$$

$$Y_1 = D_1/C_1 \tag{II}$$

$$Z_1 = Y_1 \times E_1 \tag{III}$$

$$W_1 = Y_1 \times E_1 \times F_1 \tag{IV}$$

wherein, as to the equation (I), A₁ and B₁ stand for respective half-widths of peaks, which are obtained by the process

(i) analyzing titanium oxide eight times according to an X-ray photoelectron spectroscopy,

(ii) obtaining an integrated spectrum of an electron state of titanium with respect to the above first analysis and the

(iii) obtaining a half-width, A₁, of a peak within a binding energy range of from 458 eV to 460 eV with respect to the integrated spectrum obtained in the above step (ii), and

(iv) carrying out the same steps as the above steps (ii) and (iii) with respect to the above seventh analysis and the eighth analysis to obtain a half-width, B_1 of a peak, and as to the equation (II), C_1 stands for an integrated value of absorbance within a wavelength range of from 250 nm to 550 nm in measurement of an ultraviolet-visible diffuse reflection spectrum of titanium oxide, and D_1 stands for an integrated value of absorbance of titanium oxide within a wavelength range of from 400 nm to 550 nm;

as to the equation (III), E_1 stands for a crystallite size of titanium oxide; and

as to the equation (IV), F_1 stands for a degree of anatase crystallinity thanium oxide.

The present invention also provides titanium oxide having a value of an index V_1 calculated by the following equation (V) of not more than about 0.97,

$$V_1 = H_1/G_1 \tag{V}$$

wherein, as to the equation (V), G₁ and H₁ stand for respective half-widths of peaks, which are obtained by the process

(i) analyzing titanium oxide four times according to an X-ray photoelectron spectroscopy,

(ii) obtaining an integrated spectrum of an electron state of titanium with respect to the above first analysis and the

(iii) obtaining a half-width, G₁, of a peak within a binding energy range of from 458 eV to 460 eV with respect to the

integrated spectrum obtained in the above step (ii), and

(iv) carrying out the same steps as the above steps (ii) and (iii) with respect to the above third analysis and the fourth analysis to obtain a half-width, H_1 , of a peak.



Further, the present invention provides titanium oxide having a value of an index V₁ calculated by the following equation (V) of not more than about 0.97, a value of an index U₁ calculated by the following equation (VI) of not less

> V1 = H1/G1 (V)

$$U_1 = J_1/I_1 \tag{V}$$

wherein, as to the equation (V), G_1 and H_1 stand for respective half-widths of peaks, which are obtained by the process

15

20

25

35

50

55

10

5

(i) analyzing titanium oxide four times according to an X-ray photoelectron spectroscopy,

- (ii) obtaining an integrated spectrum of an electron state of titanium with respect to the above first analysis and the
- (iii) obtaining a half-width, G₁, of a peak within a binding energy range of from 458 eV to 460 eV with respect to the
- (iv) carrying out the same steps as the above steps (ii) and (iii) with respect to the above third analysis and the fourth analysis to obtain a half-width, H_1 , of a peak; and
- as to the equation (VI), I₁ stands for an integrated value of absorbance within a wavelength range of from 220 nm to 800 nm in measurement of an ultraviolet-visible diffuse reflection spectrum of titanium oxide, and J_1 stands for an integrated value of absorbance of titanium oxide within a wavelength range of from 400 nm to 800 nm.

[0013] The present invention still further provides a photocatalyst comprising titanium oxide mentioned above as a catalyst component.

In addition, the present invention provides a photocatalytic coating agent comprising said photocatalyst and [0014] a solvent. [0015]



Further scope of applicability of the present invention will become apparent from the detailed description given hereinafter. However, it should be understood that the detailed description and specific examples, while indicating preferred embodiments of the invention, are given by way of Illustration only, since various changes and modifications within the spirit and scope of the invention will become apparent to those skilled in the art from this detailed description.

Brief Description of Drawing

[0016] The present invention will become more fully understood from the detailed description given hereinbelow and the accompanying drawings which are given by way of illustration only, and thus are not limitative of the present invention.

- Fig. 1 shows a relationship between wavelength and transmittance of an ultraviolet cut-off filter.
- Fig. 2 shows a relationship between wavelength and transmittance of an infrared cut-off filter filter.
- Fig. 3 shows a spectrum, which was obtained by the first and second analyses of titanium oxide obtained in Example 1 according to an X-ray photoelectron spectroscopy, and a spectrum, which was obtained by the seventh and 45 eighth analyses of titanium oxide obtained in Example 1 according to an X-ray photoelectron spectroscopy.
 - Fig. 4 shows a spectrum, which was obtained by the first and second analyses of titanium oxide obtained in Example 2 according to an X-ray photoelectron spectroscopy, and a spectrum, which was obtained by the seventh and eighth analyses of titanium oxide obtained in Example 2 according to an X-ray photoelectron spectroscopy.
 - Fig. 5 shows a spectrum, which was obtained by the first and second analyses of titanium oxide obtained in Comparative Example 1 according to an X-ray photoelectron spectroscopy, and a spectrum, which was obtained by the seventh and eighth analyses of titanium oxide obtained in Comparative Example 1 according to an X-ray photoelec-
 - Fig. 6 shows a spectrum, which was obtained by the first and second analyses of titanium oxide obtained in Comparative Example 2 according to an X-ray photoelectron spectroscopy, and a spectrum, which was obtained by the seventh and eighth analyses of titanium oxide obtained in Comparative Example 2 according to an X-ray photoelec-المعويد الدار



Detailed Description of the Invention

[0017] Titanium oxide (TiO_2) in accordance with the present invention has a titanium atom in a specific electron state. In order to show said electron state, the index X_1 calculated by the foregoing equation (i) is given, and its value is not more than about 0.90. The electron state can be explained by binding energy. In general, titanium oxide having a small value of the index X_1 is constituted with both a titanium atom of high binding energy electrons and a titanium atom of low binding energy electrons. When repeatedly irradiated with an X-ray, said titanium oxide is turned into titanium oxide which has a decreased titanium atom of high binding energy electrons. On the other hand, titanium oxide having a large value of the index X_1 is constituted with a titanium atom of no high binding energy electrons.

[0018] In order to obtain the integrated spectrum, titanium oxide is firstly analyzed through the following five steps in this order with use of an X-ray photoelectron spectrometer, which uses carbon as a standard for determining a peak position. Here, a time required between the beginning of the first step and the completion of the fifth step is within 30 minutes, during which titanium oxide is not exposed in the air.

1st step: analyzing an electron state of titanium two times, provided that a time per analysis is 60 seconds (the first and second analyses),

2nd step: analyzing an electron state of oxygen two times, provided that a time per analysis is 56 seconds; analyzing an electron state of carbon two times, provided that a time per analysis is 80 seconds; and analyzing an electron state of titanium two times, provided that a time per analysis is 60 seconds (the third and forth analyses),

3rd step: analyzing an electron state of oxygen two times, provided that a time per analysis is 56 seconds; analyzing an electron state of carbon two times, provided that a time per analysis is 80 seconds; and analyzing an electron state of titanium two times, provided that a time per analysis is 60 seconds (the fifth and sixth analyses),

4th step: analyzing an electron state of oxygen two times, provided that a time per analysis is 56 seconds; analyzing an electron state of carbon two times, provided that a time per analysis is 80 seconds; and analyzing an electron state of titanium two times, provided that a time per analysis is 60 seconds (the seventh and eighth analyses), and

5th step: analyzing an electron state of oxygen two times, provided that a time per analysis is 56 seconds; and analyzing an electron state of carbon two times, provided that a time per analysis is 80 seconds.

[0019] After that, X-ray photoelectron spectra are obtained as to respective analyses of from the first step to the fifth step, and thereafter, respective integrated spectra of an electron state of titanium are obtained as to the X-ray photoelectron spectrum in the first analysis and that in the second analysis. Similarly, respective integrated spectra of an electron state of titanium are obtained as to the X-ray photoelectron spectrum in the seventh analysis and that in the eighth analysis.

[0020] A half-width of a peak is obtained from a peak of titanium present within a binding energy range of from 458 eV to 460 eV in the integrated spectrum obtained above. When two or more peaks are found within the binding energy range of from 458 eV to 460 eV, a half-width of a peak is obtained from the highest peak among them.

[0021] The Index Y₁ calculated by the foregoing equation (II) shows an absorbing power of titanium oxide to visible light. A value of Y₁ is not less than about 0.075, preferably not less than about 0.110, and more preferably not less than about 0.145.

[0022] The integrated value of absorbance means an area surrounded by the horizontal axis and the diffuse reflection spectrum within the appointed wavelength range in the ultraviolet-visible diffuse reflection spectrum, provided that absorbance and wavelength are assigned to a vertical axis and a horizontal axis, respectively. The ultraviolet-visible diffuse reflection spectrum can be obtained by measurement using, for example, an ultraviolet-visible spectrophotometer and barium sulfate as a standard white board.

[0023] Among titanium oxide in accordance with the present invention, preferred is that having a value of the index Z_1 calculated by the following equation (III) of not less than about 0.75, provided that its crystallite size is assigned as E_1 .

 $Z_1 = Y_1 \times E_1 \tag{III}$

In this equation, Y_1 is an index calculated by the equation (II). More preferred value of Z_1 is not less than about 1.50, and most preferred value of Z_1 is not less than about 1.80. The crystallite size, E_1 , can be determined by obtaining a half-width of a peak and a peak position (Bragg angle) in the strongest interference line (Miller indices: 101) of titanium oxide with use of, for example, an X-ray diffractometer, and then by calculating according to Scherrer's equation.

[0024] Among titanium oxide in accordance with the present invention, preferred is that having a value of the index W_1 calculated by the following equation (IV) of not less than about 0.40, provided that a crystallite size and a degree of anatase crystallinity are assigned as E_1 and F_1 , respectively.

15

20

25



10

15

0

25

50

In this equation, Y_1 is an index calculated by the equation (II). More preferred value of W_1 is not less than about 1.30, and most preferred value of W_1 is not less than about 1.80. The degree of anatase crystallinity can be determined by obtaining a peak area in the strongest interference line (Miller indices: 101) of titanium oxide with use of, for example, [0025]

Titanium oxide (TiO₂) in accordance with the present invention has a titanium atom a specific electron state. In order to show said electron state, the index V₁ calculated by the foregoing equation (V) is given, and its value is not more than about 0.97, preferably not more than about 0.93.

In order to obtain the integrated spectrum, titanium oxide is firstly analyzed through the following second steps in this order with use of an X-ray photoelectron spectrometer, which uses carbon as a standard for determining a peak position. Here, a time required between the beginning of the first step and the completion of the second step is within 10 minutes, during which titanium oxide is not exposed in the air.

1st step : analyzing an electron state of titanium two times, provided that a time per analysis is 60 seconds (the first

2nd step : analyzing an electron state of oxygen two times, provided that a time per analysis is 56 seconds; analyzing an electron state of carbon two times, provided that a time per analysis is 80 seconds; and analyzing an electron state of titanium two times, provided that a time per analysis is 60 seconds (the third and forth analyses).

[0027] After that, X-ray photoelectron spectra are obtained as to respective analyses of from the first step to the second step, and thereafter, respective integrated spectra of an electron state of titanium are obtained as to the X-ray photoelectron spectrum in the first analysis and that in the second analysis. Similarly, respective integrated spectra of an electron state of titanium are obtained as to the X-ray photoelectron spectrum in the third analysis and that in the fourth analysis.

[0028] A half-width of a peak is obtained from a peak of titanium present within a binding energy range of from 458 eV to 460 eV in the integrated spectrum obtained above. When two or more peaks are found within the binding energy range of from 458 eV to 460 eV, a half-width of a peak is obtained from the highest peak among them.

The index U_1 calculated by the foregoing equation (VI) shows an absorbing power of titanium oxide to visible light. A value of U_1 is not less than about 0.14, preferably not less than about 0.16.

The Integrated value of absorbance means an area surrounded by the horizontal axis and the diffuse reflection spectrum within the appointed wavelength range in the ultraviolet-visible diffuse reflection spectrum, provided that absorbance and wavelength are assigned to a vertical axis and a horizontal axis, respectively. The ultraviolet-visible diffuse reflection spectrum can be obtained by measurement using, for example, an ultraviolet-visible spectrophotometer

Furthermore, titanium dioxide in accordance with the present invention has preferably anatase phase to exhibit a superior photocatalytic activity under visible light irradiation.

Shape of titanium oxide in accordance with the present invention is not particularly limited, and may be varled depending on uses (including using processes). A particle form and a fibrous form are enumerated as examples of the shape. Titanium oxide in accordance with the present invention may be used in the form of a blend with other inorganic compounds, or in the form of a composite obtainable by heating said blend, as far as the photocatalytic activity exhibited under visible light-irradiation is not impaired. Examples of said other inorganic compounds are silica (SiO₂), alumina (Al $_2$ O $_3$), zirconia (ZrO $_2$), magnesia (MgO), zinc oxide (ZnO) and iron oxide (Fe $_2$ O $_3$, Fe $_3$ O $_4$).

Titanium oxide in accordance with the present invention can be produced, for example, by a process comprising the steps of:

- (i) dissolving a commercially available titanium compound such as titanium oxysulfate, titanium tetracloride, titanium tetracloride and titanium sulfate in water to obtain a solution,
- (ii) adding a base such as ammonia, urea, an amide compound, an amidine compound and an amine compound to said solution while cooling to precipitate a solid, and
 - (iii) calcining said solid to obtain titanium oxide.

A photocatalyst in accordance with the present invention comprises titanium oxide of the present invention as a catalyst component. As the photocatalyst, there are enumerated a sheet-like photocatalyst obtained by extrusionmolding a mixture of titanium oxide in a particle form with a molding auxiliary; a sheet-like photocatalyst obtained by interweaving titanium oxide in a fibrous form with an organic fiber; and a photocatalyst obtained by coating or covering titanium oxide on a metal- or resin-made support. These photocatalysts may be used in combination with additives such as high molecular weight resins, molding auxiliaries, bonding agents, antistatic agents and absorbing agents. Alterna-



EP 1 065 169 A1

tively, these photocatalysts may be used in combination with other titanium oxide exhibiting a photocatalytic activi [0035]

With respect to a method for using the photocatalyst in accordance with the present invention, there is enu merated a method, wherein the photocatalyst and an object to be treated such as liquid to be treated and gas to b treated are placed in a glass vessel capable of transmitting visible light, and then are irradiated with visible light of wave length of not less than 430 nm. An irradiation time with visible light is not particularly limited, and may be selected approximately approxim priately depending on a strength of light in a light source and a kind and amount of a substance to be treated in th [0036]

The light source is not particularly limited, and may be capable at irradiating light containing visible light c wavelength of not less than 430 nm. Examples of the light source are sunlight, a fluorescent lamp, a halogen lamp, black light, a xenon lamp and a mercury lamp. If desired, the light source may be equipped with an ultraviolet cut-of [0037]

A photocatalytic coating agent in accordance with the present invention comprises the photocatalyst of the present invention and a solvent. The photocatalytic coating agent can be used to facilitate coating and covering of tita nium oxide on a material such as building materials and car materials, and the surface of such materials coated or cov ered with titanium oxide has a high photocatalytic activity. The solvent is not particularly limited. Preferred are those which evaporate easily after the coating or the covering, in other words, those which hardly remain on the resulting film Examples thereof are water, hydrochloric acid, alcohols and ketones.

As a process for producing the photocatalytic coating agent in accordance with the present invention, there are enumerated a process wherein titanium oxide is dispersed in water to form a slurry; and a process wherein titanium oxide is subjected to peptization with acids. When dispersing titanium oxide in the solvent, if desired, a dispersing agent [0039]

According to the present invention, there is provided titanium oxide, which exhibits a superior photocatalytic activity under not only ultrviolet light-irradiation but also visible light-irradiation. Said activity can be understood easily by a comparison between Example 1 and Comparative Example 1, or by a comparison between Example 2 and Comparative Example 2, or by a comparison between Example 3 and Comparative Example 3, or by a comparison between

The surface of building materials or car materials, which is coated with the photocatalyst or the photocatalytic coating agent in accordance with the present invention, is capable of decomposing NO_X in the air, decomposing bad-smelling substances, for example cigarette-smelling substances, present in living space and working space. decomposing undesired substances present in water such as organic solvents, agricultural chemicals and surface active agents, and further suppressing proliferation of bacteria such as radiant bacteria, algae and moulds.

Example

35

50

[0041] The present invention is illustrated in more detail with reference to the following Examples, which are only illustrative, and are not limitative for the scope of the present invention. The X-ray photoelectronic spectrophotometry spectrum of titanium oxide, the ultraviolet-visible diffuse

reflection spectrum thereof, the crystallite size thereof, the degree of anatase crystallinity thereof and the photocatalytic activity thereof were measured according to the following methods.

1. X-ray photoelectronic spectrophotometry spectrum

Measured using an apparatus of X-ray photoelectronic spectrophotometry, a trademark of XPS-7000 made by RIGAKU CORPORATION, and using the following X-ray source.

- (1) Mg was electron-irradiated at output of 8 kV and 30 mA, and the generated characteristic X-ray originated in K_{lpha} (2) Narrow scan was used as a mode of scan.
- (3) Pass E was adjusted to 10 eV.
- (4) Step E was adjusted to 0.04 eV.
- 2. Ultraviolet and visible diffuse reflection spectrum
- [0044] Measured using an ultraviolet and visible spectrophotometer, a trademark of UV-2500PC made by Shimadzu Corporation.

3. Crystallite size

Using an X-ray diffractometer, a trademark of RAD-IIA made by RIGAKU CORPORATION, a half-width, β (radian), of a peak and a peak position 2 θ (radian) in the peak of maximum intensity (Miller indices : 101) of titanium dioxide were obtained under the following conditions, followed by calculation according to the following equation (VII)

X-ray tubular bulb

:Cu

Tube voltage

:40 kV

Tube electricity Divergent slit

:35 mA :1 degree

Scattering slit

:1 degree

Light receiving slit Sampling width

:0.30 mm :0.020 degree

Scanning speed

:2.00 degree / min.

Measuring integration frequency

:1 time

20

 $E_1 (nm) = K \cdot \lambda / (\beta \cos \theta)$

(VII)

In this equation, K is a constant 0.94, and λ is a measuring X-ray wavelength (CuK $_{\alpha}$ -ray: 0.154056 nm).

4. Degree of anatase crystallinity

. 14

25

[0046] Using the same apparatus and measuring conditions as used in the above measurement of the crystallite size, a peak area in the peak of maximum intensity (Miller Indices : 101) of titanium oxide was obtained to calculate the degree of anatase crystallinity. Here, an anatase-type titanium oxide, a trademark of STT-65C-S manufactured by TITAN KOGYOU KABUSHIKI KAISHA, was used as an authentic sample, and its degree of anatase crystallinity was Example 1



In a 1-liter volume flask, 360 g of water was placed, and while stirring it, 90 g of titanium oxysulfate manufactured by SOEKAWA CHEMICAL CO., LTD. was introduced therein and mixed to obtain a solution. While cooling the solution with ice water, 101 g of an aqueous 25% ammonia solution (special grade) manufactured by Wako Pure Chemical Industries, Ltd. was added dropwise thereto over 22 minutes, thereby precipitating a solid. The solid obtained was separated by filtration and dried. The dried product obtained was calcined in air at 350 °C for 1 hour to obtain a particle titanium oxide. A crystal structure of said titanium oxide obtained was found to be an anatase-type. Physical properties of said titanium oxide and X-ray photoelectronic spectrophotometry spectrum thereof are as shown in Table 1 and Fig.

Successively, in a closed Pyrex glass-made reaction vessel having a diameter of 8 cm, a height of 10 cm and a volume of about 0.5 (iter, a glass-made petri dish having a diameter of 5 cm was placed. A photocatalyst consisting of only the particle titanium oxide obtained above was put on the petri dish. The inside of the reaction vessel was filled with a mixed gas of oxygen and nitrogen in a volume ratio of 1:4, and thereafter 4.5 µmol of acetaldehyde was put hermetically therein, followed by irradiation with visible light having wavelength of not less than 430 nm

A lighting apparatus, A trademark of OPTICAL MODULEX SX-UI500XQ made by Ushio Inc was used as a light source. The apparatus was equipped with an ultraviolet cut-off filter, a trademark of Y-45 made by ASAHI TECHNO GLASS CORPORATION, which had spectral characteristics as shown in Fig. 1, an infrared light-cutting filter, a trademark of SUPER COLD FILTER made by Ushlo Inc., which had spectral characteristics as shown in Fig. 2, and a lamp (500W xenon lamp), a trademark of UXL-500SX made by Ushlo Inc. A photocatalytic activity of the photocatalyst to actaldehyde was evaluated in terms of concentration of carbon dioxide (an oxidative decomposition product of acetaldehyde), which was generated under the irradiation, provided that said concentration was measured using a photoacoustic multi-gas monitor, 1312 type made by INNOVA. The production rate of carbon dioxide was found to be



Example 2

[0050] In a 500 mi volume eggplant-type flask, 70 g of water was placed, and while stirring it, 30 g of titanium oxysulfate manufactured by SOEKAWA CHEMICAL CO., LTD. was introduced therein and mixed to obtain a solution Using an evaporator (60 °C), the solution was concentrated to reach 86.9% by weight in terms of titanium oxysulfate While cooling the concentrated solution at -25°C in a freezer, 137 g of an aqueous 25% ammonia solution (specia grade) manufactured by Wako Pure Chemical Industries, Ltd. was added dropwise thereto over 5 seconds, thereby pre cipitating a solid. The solid obtained was separated by filtration and dried. The dried product obtained was calcined in to be an anatase-type. Physical properties of said titanium oxide. A crystal structure of said titanium oxide obtained was found to be an anatase-type. Physical properties of said titanium oxide and X-ray photoelectronic spectrophotometry spectrum thereof are as shown in Table 1 and Fig. 4, respectively.

[0051] Successively, in a manner similar to that of Example 1, a photocatalytic activity of said titanium oxide to actallytic was evaluated. The production rate of carbon dioxide was found to be 43.15 μmol/h · g-catalyst.

Comparative Example 1

[0052] β-Titanium hydroxide manufactured by KISHIDA CHEMICAL CO., LTD. was calcined in air at 400°C for 1 hour to obtain titanium oxide. Crystal structure of said titanium oxide was found to be an anatase-type. Physical properties of said titanium oxide and X-ray photoelectronic spectrophotometry spectrum thereof are as shown in Table 1 and

[0053] Successively, in a manner similar to that of Example 1, a photocatalytic activity of said titanium oxide to actaldehyde was evaluated. The production rate of carbon dioxide was found to be 0.93 µmol/h per g of catalyst.

Comparative Example 2

[0054] Example 1 was repeated, except that a photocatalyst consisting of only titanium oxide, a trademark of P-25 manufactured by DEGUSSA CORPORATION was used. The production rate of carbon dioxide was found to be 0.0 thereof are as shown in Table 1 and Fig. 6, respectively.

Example 3

25

30

In a 0.3-liter volume flask, 100 g of a 20% titanium trichloride solution (special grade) manufactured by Wako Pure Chemical was placed, and then stirred in nitrogen atmosphere. While cooling the solution with ice water, 141 g of an aqueous 25% ammonia solution (special grade) manufactured by Wako Pure Chemical Industries, Ltd. was added dropwise thereto over 30 minutes, thereby precipitating a solid. The solid obtained was separated by filtration, washed and dried. The dried product obtained was calcined in air at 400 °C for 1 hour to obtain a particle titanium oxide. A crystal structure of said titanium oxide obtained was found to be an anatase-type. Physical properties of said titanium oxide and X-ray photoelectronic spectrophotometry spectrum thereof are as shown in Table 2 and Fig. 7, respectively. Successively, in a closed Pyrex glass-made reaction vessel having a diameter of 8 cm, a height of 10 cm and a volume of about 0.5 liter, a glass-made petri dish having a diameter of 5 cm was placed. A lighting apparatus, a trademark of UI-502Q (starter: XB-50101AA-A) made by Ushio Inc., was used as a light source. The apparatus was equipped with an ultraviolet cut-off filter, a trademark of Y-45 made by ASAHI TECHNO GLASS CORPORATION, which had spectral characteristics as shown in Fig. 1 and a lamp (500W xenon lamp), a trademark of UXL-500D made by Ushlo Inc. A photocatalyst consisting of only the particle titanium oxide obtained above was put on the petri dish. The inside of the reaction vessel was filled with a mixed gas of oxygen and nitrogen in a volume ratio of 1:4, and thereafter 38 μmol of acetaldehyde was put hermetically therein, followed by Irradiation with visible light having wavelength of not less than 430 nm. A photocatalytic activity of the photocatalyst to actaldehyde was evaluated in terms of occentration of carbon dioxide (an oxidative decomposition product of acetaldehyde), which was generated under the irradiation, provided that said concentration was measured using a gas chromatograph (Shimadzu corporation). The production rate of carbon dioxide was found to be 23.4 μmol/h • g-catalyst.

Example 4

[0057] In a 0.3-liter volume flask, 25 g of a titanium tetrachloride solution (special grade) manufactured by Wako Pure Chemical Industries, Ltd. was placed, and then stirred. While cooling the solution with ice water, 36 g of an aquetos ammonia solution (special grade) manufactured by Wako Pure Chemical Industries, Ltd. was added dropwise thereto over 5 minutes, thereby precipitating a solid. The solid obtained was separated by filtration, washed and dried.

EP 1 065 169 A1

The dried product obtained was calcined in air at 400 °C for 1 hour to obtain a particle titanium oxide. A crystal structure of said titanium oxide obtained was found to be an anatase-type. Physical properties of said titanium oxide and X-ray photoelectronic spectrophotometry spectrum thereof are as shown in Table 2 and Fig. 8, respectively.

Successively, in a manner similar to that of Example 3, a photocatalytic activity of said titanium oxide to actaldehyde was evaluated. The production rate of carbon dioxide was found to be 0.75 μmol/h • g-catalyst. Comparative Example 3

[0059] Example 3 was repeated, except that a photocatalyst consisting of only titanium oxide, a trademark of P-25 manufactured by DEGUSSA CORPORATION was used. This titanium oxide had anatase and rutile phases. The production rate of carbon dioxide was found to be 0.3 μmol/h • g-catalyst. Physical properties of said titanium oxide and X-ray photoelectronic spectrophotometry spectrum thereof are as shown in Table 2 and Fig. 9, respectively.

Example 5

15

ъ

25

30

35

40

The particle titanium oxide obtained in Example 1 is dispersed in water to prepare a photocatalytic coating agent. The resulting photocatalyst coating agent is coated to car glass and then dried, and thereby it is observed that a layer of said titanium oxide is uniformly formed on the surface of car glass.

Index V / D /	Example 1	Example 2	Comparative Example 1	Comparative
Index $X_1 (=B_1/A_1)$	0.77	0.83		Example 2
Half-width A ₁ (eV)	2.20		0.86	0.99
Half-width B ₁ (eV)		1.71	1.51	1.34
ndex Y ₁ (=D ₁ /C ₁)	1.70	1.42	1.30	
	0.183	0.260	0.027	1.33
ntegrated value C ₁ of absorbance	208.3	218.9		0.065
ntegrated value D ₁ of absorbance			184.4	177.9
$ndex Z_1 (=Y_1 \times E_1)$	38.1	56.9	5.0	11.6
	3.36	3.51	0.20	
Crystallite size E1 (nm)	18.37	13.50		1.35
ndex W ₁ (=Y ₁ ×E ₁ ×F ₁)	1.11		. 7.49	20.79
Degree of anatase crystallinity F ₁ (-)		2.53	0.13	0.97
Production rate at CO (0.331	0.722	0.630	
Production rate of CO ₂ (μ mol/h • g -	19.36	43.15		0.718
atalyst)		70.10	0.93	0.0

45

50

Peak position of the integrated spectrum of first and second analysis	Example 3	Example 4	Comparative Example :
	458.7	458.2	458.2
Index V ₁ (=H ₁ /G ₁)			, 400.2
Half-width G ₁ (eV)	0.64	0.92	0.99
	2.43	1.45	1,34
Peak position of the integrated spectrum of third and fourth and sourth	458.3	458.2	458.2
Half-width H ₁ (eV)		·	100.2
ndex ⊎₁ (=J₁//₁)	1.56	1.34	1.33
4 1.17	0.196	0.119	0.126

Table 2 (continued)

rable 2 (continued)				
Integrated value I ₁ of absorbance	Example 3	Example 4	Comparative Example 3	
Integrated value J ₁ of absorbance	295.6	263.9	246.4	
Production rate of CO ₂ (μ mol/h • g -catalyst)	57.8 ·	31.4	31.0	
- g -catalyst)	23.4	0.75	0,3	
Claims			0.3	

Titanium oxide having an index X_1 calculated by the following equation (I):

$$X_1 = B_1/A_1 \tag{I}$$

of not more than about 0.90, and an index Y_1 calculated by the following equation (II):

10

15

20

25

30

35

40

45

50

55

$$Y_1 = D_1/C_1 \tag{II}$$

of not less than about 0.075, wherein A_1 and B_1 represent respective half-widths of peaks, which are obtained by the process consisting of the

- (i) analyzing titanium oxide eight times according to an X-ray photoelectron spectroscopy,
- (ii) obtaining an integrated spectrum of an electron state of titanium with respect to the above first analysis and
- (iii) obtaining a half-width, A1, of a peak within a binding energy range of from 458 eV to 460 eV with respect to the integrated spectrum obtained in the above step (li), and
- (iv) carrying out the same steps as the above steps (ii) and (ili) with respect to the above seventh analysis and the eighth analysis to obtain a half-width, B₁, of a peak,

C₁ represents the integrated value of absorbance within a wavelength range of from 250 nm to 550 nm in measurement of an ultraviolet-visible diffuse reflection spectrum of titanium oxide, and D₁ represents the integrated value of absorbance of titanium oxide within a wavelength range of from 400

2. Titanium oxide according to claim 1 having an Index Z_1 calculated by the following equation (III):

$$Z_1 = Y_1 \times E_1 \tag{III}$$

of not less than about 0.75, wherein Y_1 is as defined in claim 1 and E_1 represents the crystallite size of titanium oxide.

Titanium oxide according to claim 2 having an index W₁ calculated by the following equation (IV):

$$W_1 = Y_1 \times E_1 \times F_1 \tag{IV}$$

of not less than about 0.40,

wherein Y_1 is as defined in claim 1, E_1 is as defined in claim 2, and F_1 represents the degree of anatase crystallinity

Titanium oxide having an index V_1 calculated by the following equation (V):

$$V_1 = H_1/G_1 \tag{V}$$

of not more than about 0.97,

wherein G_1 and H_1 represent respective half-widths of peaks, which are obtained by the process consisting of the

EP 1 065 169 A1

- (i) analyzing titanium oxide four times according to an X-ray photoelectron spectroscopy,
- (ii) obtaining an integrated spectrum of an electron state of titanium with respect to the above first analysis and
- (iii) obtaining a half-width, G₁, of a peak within a binding energy range of from 458 eV to 460 eV with respect to the integrated spectrum obtained in the above step (ii), and
- (iv) carrying out the same steps as the above steps (ii) and (iii) with respect to the above third analysis and the fourth analysis to obtain a half-width, H₁, of a peak.
- Titanium oxide according to claim 4 having an index U₁ calculated by the following equation (VI):

$$U_1 = J_1/I_1 \tag{VI}$$

of not less than about 0.14,

5

10

15

30

35

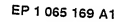
40

45

wherein l_1 represents the integrated value of absorbance within a wavelength range of from 220 nm to 800 nm in measurement of an ultraviolet-visible diffuse reflection spectrum of titanium oxide, and

 ${\sf J_1}$ represents the integrated value of absorbance of titanium oxide within a wavelength range of from 400 nm to 800 nm.

- Titanlum oxide according to claim 4, wherein said titanium has anatase phase. 50
 - Titanium oxide according to claim 5, wherein said titanium has anatase phase. 7.
- A photocatalyst comprising titanium oxide as claimed in any one of claims 1 to 7 as a catalyst component. 8. 25
 - A photocatalytic coating agent, which comprises:
 - (i) a soivent; and
 - (II) a photocatalyst comprising titanium oxide as claimed in any one of claims 1 to 7 as a catalyst component.
 - 10. A material coated with a photocatalyst as claimed in claim 8 or photocatalytic coating agent as claimed in claim 9.
 - 11. Use of titanium oxide as claimed in any one of claims 1 to 7 as a photocatalyst.



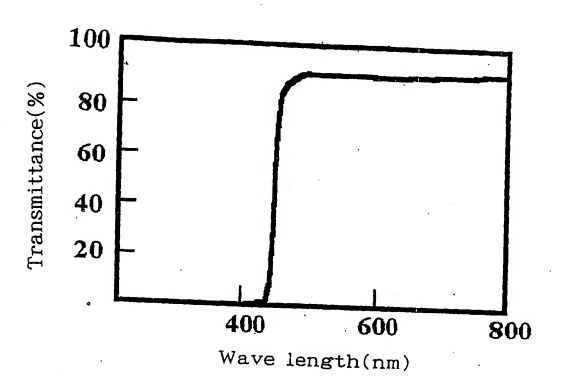


Fig.1

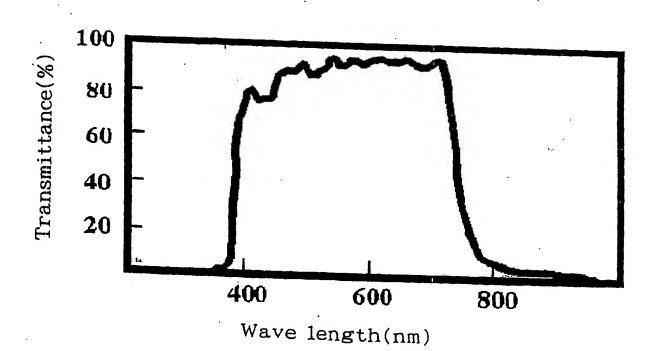
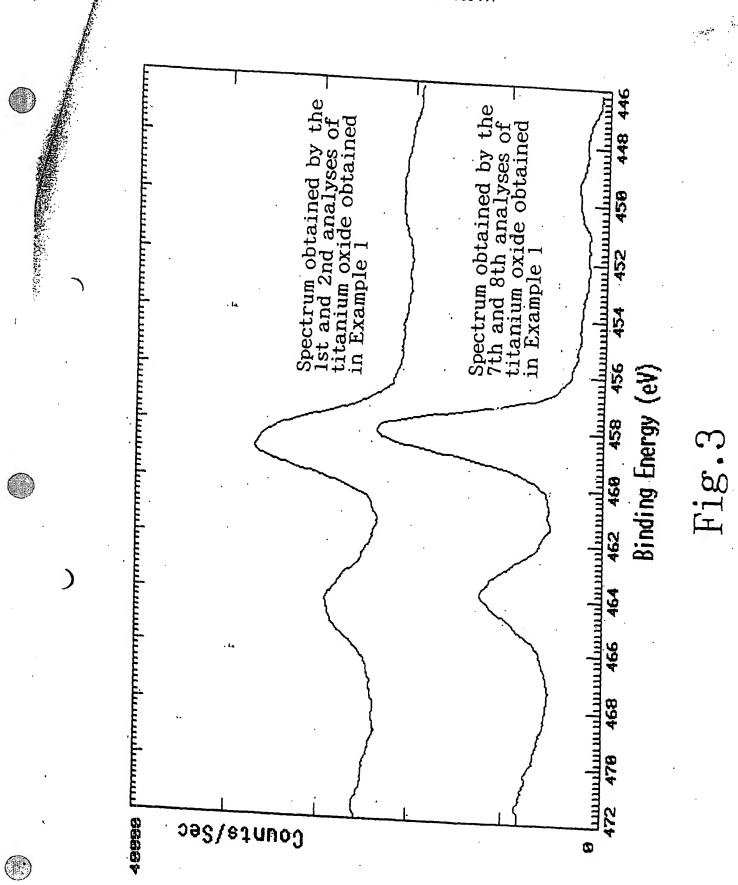
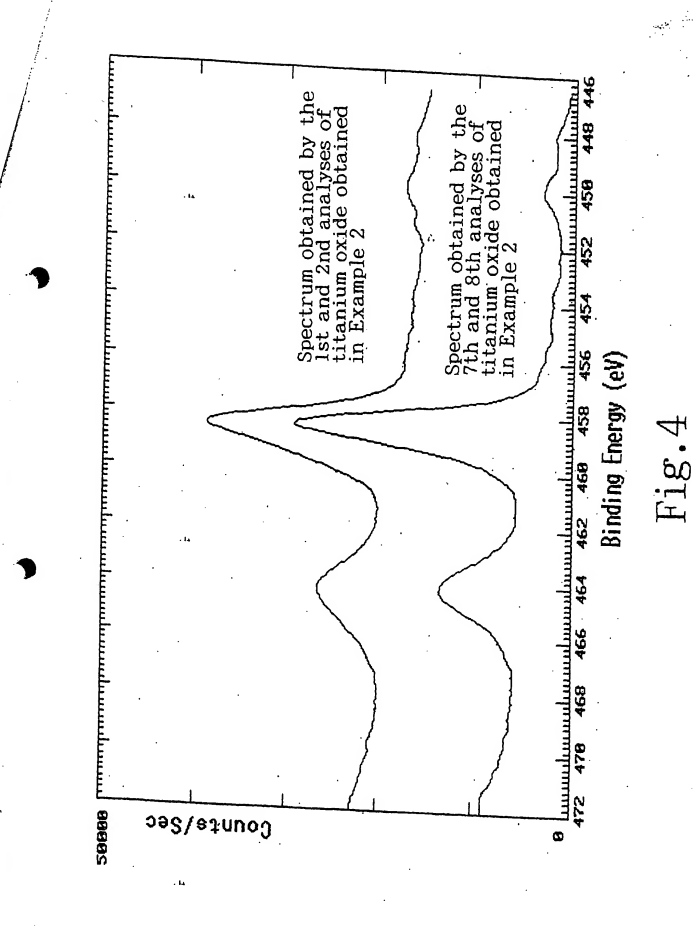
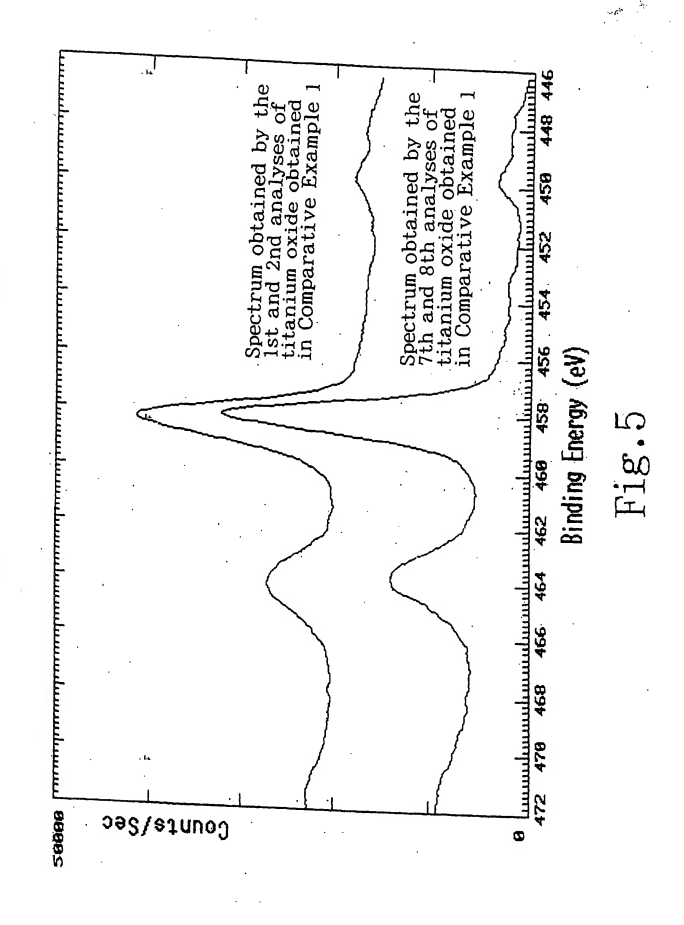
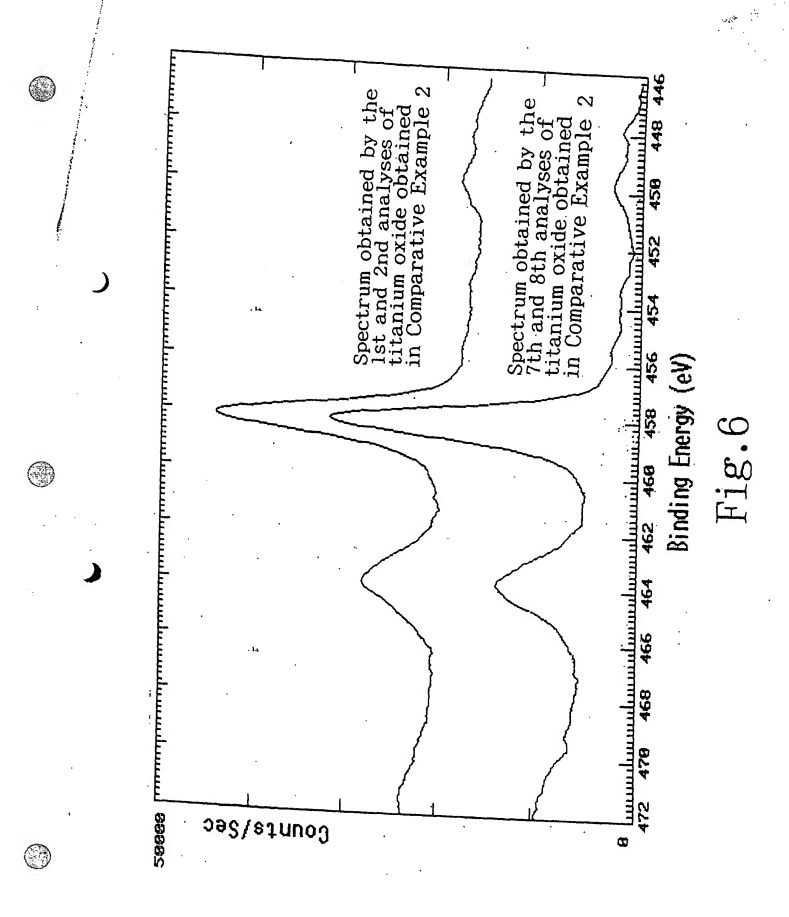


Fig.2











EUROPEAN SEARCH REPORT

Application Number

EP 00 30 5433

Calcas	Citation of de-	IDERED TO BE RELEVANT		
Category	of relevant p	th indication, where appropriate, * assages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CI.7)
A	DE 34 39 217 A (K 30 April 1986 (19	RONOS TITAN GMBH) 86-04-30)		C01G23/047 C01G23/053
A	EP 0 923 988 A (N 23 June 1999 (199	IPPON SODA CO) 9-06-23)	·	B01J21/06 B01J35/00
4	EP 0 675 086 A (I 4 October 1995 (1	SHIHARA SANGYO KAISHA) 995-10-04)		
	DATABASE WPI Section Ch, Week Derwent Publication Class D21, AN 1990 XP002148842 å JP 04 367512 A (18 December 1992 (* abstract *	ons Ltd., London, GB; 3-041323		
	·			TECHNICAL FIELDS SEARCHED (Int.CI.7)
				C01G B01J
	. l.a	•		
				•
				·
1	he present search report has	been drawn up for all claims	1	
P	lace of search	Date of completion of the search		
	HE HAGUE	29 September 200	0 Thior	Examiner n M
X : particul Y : particul docume A : technol O : non-wi	any relevant it taken alone atry relevant it taken alone atry relevant it combined with and of the same category operations background than disclosure diate document	T : theory or principl E : earlier patent doc after the filling dat ther D : document cited is L : document cited for	e underlying the involunt to underlying the involunt, but published to the application of either reasons	ention ed on, or

EPO FORM 1503 03.82 (POJCO1)

ANNEX TO THE EUROPEAN SEARCH REPORT ON EUROPEAN PATENT APPLICATION NO.

EP 00 30 5433

This annex lists the patent family members relating to the patent documents cited in the above-mentioned European search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

29-09-2000

Patont do					29-09-2
Patent document cited in search report		Publication date		Patent family member(s)	Publication
DE 3439217	Α	30-04-1986	NON	<u> </u>	date
EP 0923988 EP 0675086	Α	23-06-1999	JP JP WO	9310039 A 3038599 B 9700134 A	02-12-199 08-05-200 03-01-199
. 14		04-10-1995	AT AU CA CN DE DX EP SG JP JP	147366 T 4216893 A 2101360 A 1093060 A,B 1159466 A 69307208 D 69307208 T 581216 T 0581216 A 2096152 T 6293519 A 42893 A 7002522 A 7000819 A	15-01-1997 03-02-1994 29-01-1994 17-09-1997 20-02-1997 05-06-1997 02-02-1994 01-03-1997 21-10-1994 17-10-1995 06-01-1995
JP 4367512	A	18-12-1992	JP	2709206 B	04-02-1998

For more details about this annex : see Official Journal of the European Patent Office, No. 12/82